

Graph Convolution Networks

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Outline

1 Graph Convolutional Networks

- Why GCN
- How to extend convolution to graphs?

2 Spatial Approach

3 Spectral Approach

- Basics of Spectral Approach
- Problem Formulation
- Graph Laplacian

4 Spectral Networks and Deep Locally Connected Networks on Graphs

5 CNN on Graphs with Fast Localized Spectral Filtering

- Learning fast localized Spectral filters
- Coarsening and Pooling
- Results on MNIST

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Introduction to Graph Convolutional Networks

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- But CNN donot directly generalize to irregular domains such as graph.
- Want to generalize CNN to Graphs.
- Non-trivial because the distances are non-euclidean.

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Extending Convolutional to Graphs

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Generalization of CNN in the spatial domain itself.

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Using the frequency characterization of CNN and using that to generalize to Graphical domain

Extending Convolutional to Graphs

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- Spatial Approach :
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- Spectral Approach :
Using the frequency characterization of CNN and using that to generalize to Graphical domain
 - ▶ Spectral Networks and Deep Locally Connected Networks on Graphs [Bruna et al. ICLR 2014].
 - ▶ Convolutional Neural Networks on Graphs with Fast Localized Spectral Filtering [Defferrard et al. NIPS 2016] (will be the main focus)
 - ▶ Semi-Supervised Classification with Graph Convolutional Networks [Kipf et al. ICLR 2017]

Limitations of Spatial Approach

- Can't exactly define a neighborhood because the distances are not uniform.
- Ordering of nodes is problem specific.

Hence for the remainder we discuss the Spectral Approach

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A Basic Formulation

- Convolution in spectral (Fourier) domain is point wise multiplication.
- Fourier Basis is defined as the eigen basis of the laplacian operator.
- Can use Laplacian of a graph.

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Defining the Problem on Graphs

- A feature description x_i for every node i ; summarized in a $N \times D$ feature matrix X (N : number of nodes, D : number of input features)

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- Adjacency Matrix A .
- Node level output Z (an $N \times F$ feature matrix, where F = number of output features per node).

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Brief overview of Graph Laplacian

Let T denote the diagonal matrix with (v,v) -th entry having value d_v : degree of vertex v . Define L-matrix as

$$L(u, v) = \begin{cases} d_v & \text{if } u = v \\ -1 & \text{if } u \text{ and } v \text{ are adjacent} \\ 0 & \text{otherwise} \end{cases}$$

And the Laplacian of the graph as

$$\mathcal{L}(u, v) = \begin{cases} 1 & \text{if } u = v \\ -\frac{1}{\sqrt{d_u d_v}} & \text{if } u \text{ and } v \text{ are adjacent} \\ 0 & \text{otherwise} \end{cases}$$

Graph Laplacian (contd.)

$$\mathcal{L} = T^{-1/2} L T^{1/2}$$

With the convention $T^{-1}(v, v) = 0$ for $d_v = 0$.

When G is k -regular,

$$\mathcal{L} = I - \frac{1}{k} A$$

For a general graph

$$\mathcal{L} = I - T^{-1/2} A T^{1/2}$$

Spectral Network Approach

- Mentions the use of both spatial and spectral construction.
- For the spectral part uses a spline and has k control points for it.

$$g_{\theta}(\Lambda) = B\theta$$

Here B is the cubic B-spline basis and θ is a vector of control points.

- The datasets used (created) are quite interesting. Subsampled MNIST and MNIST on sphere to show how spectral networks can be used on graphs.

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Graph Fourier Transform

- Laplacian of the graph is real symmetric positive semidefinite, and thus can be written as

$$L = U\Lambda U^T$$

- Here $U = [u_0 \dots u_{n-1}]$ is the fourier basis and $\Lambda = \text{diag}([\lambda_0 \dots \lambda_{n-1}])$ are ordered real non-negative eigen values.
- Graph Fourier Transform of a signal x is $\hat{x} = U^T x$.

Spectral filtering of graph signals

- Defining convolution on graphs

$$x *_G y = U((U^T x) \odot (U^T y))$$

- Filtering by g_θ

$$y = g_\theta(L)x = g_\theta(U\Lambda U^T)x = Ug_\theta(\Lambda)U^T x$$

- A non-parametric filter (all parameters free) would be defined as

$$g_\theta(\Lambda) = \text{diag}(\theta)$$

Polynomial Parametrization

- Problem with non-parametric filters is that not localized (we want something like k-neighborhood) and therefore their learning complexity becomes $O(n)$. This can be overcome with use of a Polynomial filter

$$g_{\theta}(\Lambda) = \sum_{k=0}^{K-1} \theta_k \Lambda^k$$

- The advantage we gain here is that nodes which are at a distance greater than K away from the node i , at which the filter is applied, are not affected. Hence we have gained localization.

Recursive formulation for fast filtering

- Still cost to filter is high $O(n^2)$ because of multiplication with U matrix.
- Therefore use recurrence relation of chebyshev polynomial instead.

$$g_{\theta}(\Lambda) = \sum_{k=0}^{K-1} \theta_k T_K(\tilde{\Lambda})$$

Here $\tilde{\Lambda}$ is scaled between $[-1, 1]$.

- This allows us to compute $\bar{x}_k = T_K \tilde{L}x$. And Therefore

$$y = g_{\theta}(L)x = [\bar{x}_0 \dots \bar{x}_{k-1}] \theta$$

- The cost is now $O(K|E|)$

Learning filters

- Trivial to show that backprop calculation can be done efficiently.

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Graph Coarsening and Pooling

- Require efficient mechanism for pooling. Graph clustering as such is NP-hard and some approximations must be made.
- The paper uses Graclus algorithm for coarsening, and uses an intelligent way of rearranging the nodes [creating a balanced binary tree from the remaining singleton and fake nodes] so that the pooling now becomes equivalent to pooling a regular 1D signal.

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MNIST results

- Achieves close to classical CNN accuracy.
- Pictures to be added.



M. Niepert, M. Ahmed, and K. Kutzkov, “Learning convolutional neural networks for graphs,” *CoRR*, vol. abs/1605.05273, 2016.